

STIC Database Tracking Number: 108474

TO: Devesh Khare

Location: cm1/8a13/8b19

Art Unit: 1623

Tuesday, November 18, 2003

Case Serial Number: 10/007489

From: Susan Hanley

Location: Biotech-Chem Library

CM1 6B05

Phone: 305-4053

susan.hanley@uspto.gov

Search Notes			
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TIC TOB Search Request Form

108474 WStreet War 127/http://ptoweb/patents/stic/searchsubmit

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Tech Center:				
○ TC 1600○ TC 1700○ TC 2100○ TC 2600○ TC 2800○ TC 3600○ TC 3700○ Other				
Enter your Contact Information below:				
Name:				
Devesh Khare				
Employee Number: 77931 Phone:				
605-1199				
Art Unit or Office: 1623 Building & Room Number:				
8 A 13, Mail 8 B19				
Enter the case serial number (Required): 10/007,489				
If not related to a patent application, please enter NA here.				
Class / Subclass(es) 536/25.34				
Earliest Priority Filing Date: 09/14/1998				
Format preferred for results: Paper Diskette E-mail				

Provide detailed information on your search topic:

- In your own words, describe in detail the concepts or subjects you want us to search.
- Include synonyms, keywords, and acronyms. Define terms that have special meanings.
- *For Chemical Structure Searches Only*
 Include the elected species or structures, keywords, synonyms, acronyms, and registry numbers

- *For Sequence Searches Only*
 Include all pertinent information (parent, child, divisional, or issued patent numbers) along with the appropriate serial number.
- *For Foreign Patent Family Searches Only* Include the country name and patent number.
- Provide examples or give us relevant citations, authors, etc., if known.
- FAX or send the abstract, pertinent claims (not all of the claims), drawings, or chemical structures to your EIC or branch library.

Please search the following claims:	:	
Claim 1: A method for generating phosphorot comprising: 1) growing a single-stranded recombinant DN that uses thio-phosphate as a source of pho	A phage in m	
2) harvesting the single-stranded phage and corresponding to the recombinant DNA insert	purifying th	e DNA
3) fragmentation of the insert DNAsuch that	oligo mixtu	res spanning
the entire length of the segment are general	ited	
the entire length of the segment are general Claim 2: the method of claim 1 used to general DNA, ss DNA, and/or RNA by in vivo incorpo	erate phospho	rothicate ds o-phosphate
the entire length of the segment are general claim 2: the method of claim 1 used to general DNA, ss DNA, and/or RNA by in vivo incorpo	erate phospho	rothioate ds o-phosphate
the entire length of the segment are general claim 2: the method of claim 1 used to general control of the segment are general control of the	erate phospho	rothioate ds o-phosphate
the entire length of the segment are general claim 2: the method of claim 1 used to general conditions of the segment are general conditions. The condition is a segment are general conditions of the conditions	erate phospho	rothicate ds o-phosphate
the entire length of the segment are general claim 2: the method of claim 1 used to general control of the segment are general control of the	erate phospho	rothioate ds o-phosphate
the entire length of the segment are general claim 2: the method of claim 1 used to general DNA, as DNA, and/or RNA by in vivo incorposinto nucleotide precursor pools. Thank you.	erate phospho	rothicate ds o-phosphate
the entire length of the segment are general the entire length of the segment are general than 2: the method of claim 1 used to general than 2: the method of claim 1 used to general than 3: the method of claim 1 used that 3: the method of claim 1 used that 3: the method of claim 1 used that 3: the method of claim	erate phospho	rothioate ds o-phosphate

Press ALT + F, then P to print this screen for your own information.

SEND	DESET
SEND	RESET

USPTO <u>întranet Home</u> | <u>Index</u> | <u>What's New</u> | <u>Resources</u> | <u>Contacts</u> | <u>Internet</u> | <u>Search</u> | <u>Firewall</u> | <u>Web</u> Services

Last Modified: Wednesday, December 31, 1969 19:00:00

-> file medline FILE 'MEDLINE' ENTERED AT 14:47:28 ON 18 NOV 2003

FILE LAST UPDATED: 13 NOV 2003 (20031113/UP). FILE COVERS 1958 TO DATE.

On April 13, 2003, MEDLINE was reloaded. See HELP RLOAD for details.

MEDLINE thesauri in the /CN, /CT, and /MN fields incorporate the MeSH 2003 vocabulary. See http://www.nlm.nih.gov/mesh/changes2003.html for a description on changes.

This file contains CAS Registry Numbers for easy and accurate substance identification.

-> d que 158

LSS	* * * * * * * * * * * * * * * * * * * *	PLU=0N	10101-88-9 OR THIOPHOSPHORIC
	OR 13598-51-1	D D.	1.55 AND (CCDM) OD CTMCLE CTDAM
LS6	3 SEA FILE=MEDLINE ABB=ON D? OR SS DNA)	PLU=UN	L55 AND (SSDNA OR SINGLE-STRAN
LS7		PI II-ON	ORGANOTHIOPHOSPHORUS COMPOUNDS
C3,	/CT	7 40-011	
L58	1 SEA FILE-MEDLINE ABB-ON	PLU=ON	LS6 AND LS7

-> file embase

FILE 'EMBASE' ENTERED AT 14:47:29 ON 18 NOV 2003 COPYRIGHT (C) 2003 Elsevier Inc. All rights reserved.

FILE COVERS 1974 TO 13 Nov 2003 (20031113/ED)

EMBASE has been reloaded. Enter HELP RLOAD for details.

This file contains CAS Registry Numbers for easy and accurate substance identification.

=> d que 173

L64 962 SEA FILE=EMBASE ABB=ON	PLU=ON 10101-88-9 OR THIOPHOSPHORIC
OR 13598-51-1 OR THIOPH	OSPHATE
L65 69 SEA FILE=EMBASE ABB=ON	PLU=ON L64 AND (SSDNA OR SINGLE-STRAND
? OR SS DNA OR ?PHAGE O	R PLASMID)
L66 38 SEA FILE=EMBASE ABB=ON	PLU=ON L6S AND ?OLIGO?
L70 15 SEA FILE=EMBASE ABB=ON	PLU=ON L66 AND (HIGH OR THIOPHOSPHATE
OR PHOSPHOROTHIOATE OR	EXTENDING)/TI
L71 3 SEA FILE=EMBASE ABB=ON	PLU=ON L70 NOT (CHIRAL OR EFFECT OR
VIRAL OR VIVO OR ANTIP	ARALLEL OR SFII OR MICE OR MACROPHAGE
OR GENE OR CPG)/TI	
L72 1 SEA FILE=EMBASE ABB=ON	PLU=ON L66 AND EXTENDING/TI
L73 4 SEA FILE=EMBASE ABB=ON	

⇒ file hcaplus

FILE 'HCAPLUS' ENTERED AT 14:47:30 ON 18 NOV 2003
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FILE COVERS 1907 - 18 Nov 2003 VOL 139 ISS 21 FILE LAST UPDATED: 17 Nov 2003 (20031117/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

```
-> d que 110
           1813) SEA FILE-HCAPLUS ABB-ON PLU-ON PHOSPHOROTHIOATE OLIGONUCLEOTI
L1 (
            DES+PFT,NT/CT
231 SEA FILE-HCAPLUS ABB=ON PLU=ON L1(L)PREP/RL
12
              5 SEA FILE-REGISTRY ABB-ON PLU-ON 03PS/MF
L4
           8754 SEA FILE-REGISTRY ABB=ON PLU=ON
                                                    "PHOSPHOROTHICATE"
L5
            448 SEA FILE-REGISTRY ABB=ON
                                           PLU=ON
                                                   LS AND M/ELS
L6
             35 SEA FILE=REGISTRY ABB=ON
                                           PLU=ON
                                                   L6 NOT C/ELS
.L7
             40 SEA FILE=REGISTRY ABB=ON PLU=ON L4 OR L7
18
            354 SEA FILE-HCAPLUS ABB=ON PLU=ON L8
L9
              2 SEA FILE=HCAPLUS ABB=ON PLU=ON L9 AND L2
L10
=> d que 119
           1813 SEA FILE-HCAPLUS ABB=ON PLU=ON PHOSPHOROTHIOATE OLIGONUCLEOTI
1.3
                DES+PFT.NT/CT
              5 SEA FILE=REGISTRY ABB=ON PLU=ON 03PS/MF
L4
           8754 SEA FILE=REGISTRY ABB=ON
                                           PLU=ON
                                                    "PHOSPHOROTHIOATE"
L5
            448 SEA FILE-REGISTRY ABB=ON
                                            PLU=0N
                                                    LS AND M/ELS
L6
             35 SEA FILE=REGISTRY ABB=ON
                                            PLU=ON
                                                    L6 NOT C/ELS
L7
             40 SEA FILE=REGISTRY ABB=ON
                                           PLU=ON
                                                   L4 OR L7
L8
            354 SEA FILE-HCAPLUS ABB=ON
                                          PLU⇒ON
                                                   L8
L9
         222353 SEA FILE=HCAPLUS ABB=ON
                                          PLU=ON
                                                   DNA+PFT/CT
L16
           5268 SEA FILE=HCAPLUS ABB=ON
                                          PLU=0N
                                                   L16(L)(SS OR SINGLE-STRAND?)
L17
             24 SEA FILE-HCAPLUS ABB=ON
                                          PLU=ON
                                                   L17 AND L3
L18
L19
              1 SEA FILE=HCAPLUS ABB=ON
                                          PLU=ON
                                                   L9 AND L18
=> d que 123
           1813 SEA FILE=HCAPLUS ABB=ON PLU=ON PHOSPHOROTHIOATE OLIGONUCLEOTI
L3
                DES+PFT, NT/CT
                                         PLU=ON DNA+PFT/CT
PLU=ON L16(L)(SS (
L16
         222353 SEA FILE-HCAPLUS ABB=ON
                                                   L16(L)(SS OR SINGLE-STRAND?)
L17
           5268 SEA FILE-HCAPLUS ABB=ON
                                          PLU=ON
L18
             24 SEA FILE-HCAPLUS ABB=ON
                                                   L17 AND L3
                                                   (MONOTHIO? OR PHOSPHOROTHIO?
                                          PLU=ON
L20
         486562 SEA FILE=HCAPLUS ABB=ON
                OR THIO? OR PHOSPHOROMONOTHIO?)
L21
             24 SEA FILE-HCAPLUS ABB-ON
                                          PLU=ON
                                                   L20 AND L18
                                                  L21 AND (PHAGE OR BACTERIOPHAG
              3 SEA FILE-HCAPLUS ABB=ON
                                          PLU=ON
L22
              2 SEA FILE-HCAPLUS ABB=ON PLU=ON L22 NOT CIRCULAR/TI
L23
-> d que 125
           1813) SEA FILE-HCAPLUS ABB-ON PLU-ON PHOSPHOROTHIOATE OLIGONUCLEOTI
L1 (
         DES+PFT,NT/CT
231 SEA FILE=HCAPLUS ABB=ON
222353 SEA FILE=HCAPLUS ABB=ON
                                          PLU=ON
                                                   L1(L)PREP/RL
12
L16
                                          PLU=ON
                                                   DNA+PFT/CT
           5268 SEA FILE-HCAPLUS ABB-ON
                                          PLU=ON
                                                   L16(L)(SS OR SINGLE-STRAND?)
L17
             12 SEA FILE=HCAPLUS ABB=ON
                                          PLU=ON
                                                   L2 AND (PHAGE OR BACTERIOPHAGE
L24
L25
              2 SEA FILE-HCAPLUS ABB=ON PLU=ON L24 AND L17
=> d que 141
              5 SEA FILE~REGISTRY ABB=ON
                                           PLU=ON 03PS/MF
L5
           8754 SEA FILE=REGISTRY ABB=ON
                                            PLU=0N
                                                    "PHOSPHOROTHIOATE"
            448 SEA FILE-REGISTRY ABB=ON
                                            PLU=ON
                                                    L5 AND M/ELS
L7
             35 SEA FILE=REGISTRY AB8=ON
                                           PLU=ON
                                                    L6 NOT C/ELS
             40 SEA FILE=REGISTRY ABB=ON
                                          PLU=ON
                                                   L4 08 L7
L9
            354 SEA FILE=HCAPLUS ABB=ON PLU=ON
                                                   L8
                                                   L9 AND L5
L32
            354 SEA FILE-HCAPLUS ABB-ON
                                          PLU=ON
                                                  L32 AND SINGLE-STRAND?
L33 AND POLICOP
L39 AND PTHIOP
                                          PLU=ON
L33
             11 SEA FILE=HCAPLUS ABB=ON
L39
             10 SEA FILE-HCAPLUS ABB=ON
                                          PLU=ON
                                          PLU=ON
             10 SEA FILE=HCAPLUS ABB=ON
L40
                                                   L40 NOT (GOLD OR DOUBLE OR
                SEA FILE-HCAPLUS ABB=ON
                                          PLU=ON
L41
                HAPLOTYPES)/TI
=> s 110 or 119 or 123 or 125 or 141
            12 L10 OR L19 OR L23 OR L25 OR L41
L74
=> dup rem 158 173 174
```

```
FILE 'MEDLINE' ENTERED AT 14:47:58 ON 18 NOV 2003
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PROCESSING COMPLETED FOR LS8
PROCESSING COMPLETED FOR L73
PROCESSING COMPLETED FOR L74
                17 DUP REM LS8 L73 L74 (O DUPLICATES REMOVED)
ANSWER '1' FROM FILE MEDLINE
ANSWERS '2-5' FROM FILE EMBASE
ANSWERS '6-17' FROM FILE HCAPLUS
175
=> d ibib abs ind 1-5
     ANSWER 1 OF 17
                               MEDLINE on $TN
ACCESSION NUMBER:
                          85054878
                                           MEDLINE
DOCUMENT NUMBER:
                          85054878
                                         PubMed ID: 6094546
TITLE:
                          Cleavage of phosphorothioate-substituted DNA by restriction
                          endonucleases.
AUTHOR:
                          Potter B V; Eckstein F
SOURCE:
                          JOURNAL OF BIOLOGICAL CHEMISTRY, (1984 Nov 25) 259 (22)
                          14243-8.
                          Journal code: 2985121R. ISSN: 0021-9258.
PUB. COUNTRY:
                          United States
DOCUMENT TYPE:
                          Journal; Article; (JOURNAL ARTICLE)
LANGUAGE:
                          English
FILE SEGMENT:
                          Priority Journals
ENTRY MONTH:
                          198412
ENTRY DATE:
                          Entered STN: 19900320
                          Last Updated on STN: 19900320
                          Entered Medline: 19841227
      M13 RF DNA was synthesized in vitro in the presence of various single deoxynucleoside 5'-O-(1-thiotriphosphate) phosphorothioate analogues, and the three other appropriate deoxynucleoside triphosphates using a M13 (+)-
AB
      single-stranded template, Escherichia coli DNA
polymerase I and T4 DNA ligase. The resulting DNAs contained various
restriction endonuclease recognition sequences which had been modified at
      their cleavage points in the (-)-strand by phosphorothioate substitution. The behavior of the restriction enzymes Aval, BamHI, EcoRI, HindIII, and
       Sall towards these substituted DNAs was investigated. EcoRI, BamHI, and
      HindIII were found to cleave appropriate phosphorothioate-substituted DNA
      at a reduced rate compared to normal M13 RF DNA, and by a two-step process
       in which all of the DNA is converted to an isolable intermediate nicked
      molecule containing a specific discontinuity at the respective recognition
      site presumably in the (+)-strand. By contrast, SalI cleaved substituted
      DNA effectively without the intermediacy of a nicked form. Aval, however,
       is only capable of cleaving the unsubstituted (+)-strand in appropriately
      modified DNA.
      Check Tags: Support, Non-U.S. Gov't
Bacteriophage phi X 174: GE, genetics
        Base Sequence
        Binding Sites
       *DNA Restriction Enzymes: ME, metabolism
         *DNA, Single-Stranded: AN, analysis
        DNA, Viral: AN, analysis
        Deoxyribonuclease BamHI
        Deoxyribonuclease EcoRI
        Deoxyribonuclease HindIII
         *Organothiophosphorus Compounds: ME, metabolism
         *Thiophosphoric Acid Esters: ME, metabolism
      0 (DNA, Single-Stranded); 0 (DNA, Viral); 0 (Organothiophosphorus Compounds); 0 (Thiophosphoric Acid
      Esters); EC 3.1.21 (DNA Restriction Enzymes); EC 3.1.21.- (Deoxyribonuclease BamHI); EC 3.1.21.- (Deoxyribonuclease EcoRI); EC 3.1.21.- (Deoxyribonuclease HindIII); EC 3.1.21.- (endodeoxyribonuclease AvaI); EC 3.1.21.- (endodeoxyribonuclease SalI)
L75 ANSWER 2 OF 17 EMBASE COPYRIGHT 2003 ELSEVIER INC. ALL RIGHTS RESERVED.
      on STN
ACCESSION NUMBER:
                          96053473 EMBASE
DOCUMENT NUMBER:
                          1996053473
TITLE:
                          Extending the chemistry that supports genetic
```

information transfer in vivo: Phosphorothicate DNA,

```
phosphorothioate RNA, 2'-0-methyl RNA, and
                              methylphosphonate DNA.
                              Thaler D.S.; Liu S.; Tombline G.
DNA RMCP, Jefferson Cancer Center, Thomas Jefferson
AUTHOR:
CORPORATE SOURCE:
                              University, 233 South 10th Street, Philadelphia, PA 19107.
                              United States
                              Proceedings of the National Academy of Sciences of the United States of America, (1996) 93/3 (1352-1356). ISSN: 0027-8424 CODEN: PNASA6
SOURCE:
COUNTRY:
                              United States
DOCUMENT TYPE:
                              Journal; Article
FILE SEGMENT:
                              022
                                          Human Genetics
                              029
                                          Clinical Biochemistry
LANGUAGE:
                              English
SUMMARY LANGUAGE:
                              English
       DNA and RNA are the polynucleotides known to carry genetic information in
       life. Chemical variants of DNA and RNA backbones have been used in structure- function and biosynthesis studies in vitro, and in antisense
       pharmacology, where their properties of nuclease resistance and enhanced cellular uptake are important. This study addressed the question of whether the base(s) attached to artificial backbones encodes genetic information that can be transferred in vivo. Oligonucleotides
       containing chemical variants of DNA or RNA were used as primers for site-specific mutagenesis of bacteriophage fl. Progeny phage were scored both genetically and physically for the inheritance of information originally encoded by bases attached to the
       nonstandard backbones. Four artificial backbone chemistries were tested: phosphorothioate DNA, phosphorothioate RNA, 2'-0-methyl RNA and methylphosphonate DNA. All four were found capable of faithful information transfer from their attached bases when one or three artificial positions
       were flanked by normal DNA. Among oligonucleotides composed entirely of nonstandard backbones, only phosphorothioate DNA supported
       genetic information transfer in vivo.
Medical Descriptors:
        *gene transfer
       *nucleotide sequence
       article
       chemical structure
       dna replication
       dna synthesis
       genetic code
       molecular genetics
       priority journal site directed mutagenesis
       structure activity relation
       Drug Descriptors:
       *dna
       *rna
          antisense oligonucleotide
          oligonucleotide
       phosphorothioic acid
        transfer rna
       (dna) 9007-49-2; (rna) 63231-63-0; (phosphorothioic acid)
       10101-88-9, 13598-51-1, 15181-41-6; (transfer rna)
       9014-25-9
L75 ANSWER 3 OF 17 EMBASE COPYRIGHT 2003 ELSEVIER INC. ALL RIGHTS RESERVED.
       on STN
ACCESSION NUMBER:
                              93173385 EMBASE
DOCUMENT NUMBER:
                              1993173385
TITLE:
                              Site-directed mutagenesis of single-
                              stranded and double-stranded DNA by
                              phosphorothicate approach.
                              Olsen D.B.; Sayers J.R.; Eckstein F.
Methods in Enzymology, (1993) 217/- (189-217).
ISSN: 0076-6879 CODEN: MENZAU
AUTHOR .
SOURCE:
                              United States
COUNTRY:
                              Journal; Article
029 Clinical Biochemistry
DOCUMENT TYPE:
FILE SEGMENT:
LANGUAGE:
                              English
       Medical Descriptors:
       *site directed mutagenesis
       article
          bacteriophage t7
       cell transformation
       dna sequence
       dna synthesis
       dna template
```

```
escherichia coli
      gene mutation
      hydrolysis
      nonhuman
      nucleotide sequence
        plasmid
      polymerization
      priority journal
      Drug Descriptors:
      *double stranded dna
      *phosphorothioic acid
         *plasmid dna
         *single stranded dna
      dna polymerase
      ethidium bromide
      exodeoxyribonuclease iii
        oligonucleotide
      primer dna
      restriction endonuclease
      (phosphorothioic acid) 10101-88-9, 13598-51-1,
      15181-41-6; (dna polymerase) 37217-33-7; (ethidium bromide) 1239-45-8;
      (exodeoxyribonuclease iii) 9037-44-9
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      on STN
ACCESSION NUMBER:
                        90080670 EMBASE
                         1990080670
DOCUMENT NUMBER:
TITLE:
                        High-efficiency oligonucleotide
                         -directed plasmid mutagenesis.
AUTHOR:
                        Olsen D.B.; Eckstein F.
                        Max-Plank Institut fur, Experimentelle Medizin, Abteilung
Chemie, Hermann-Rein Strasse 3,D-3400 Cottingen, Germany
CORPORATE SOURCE:
                        Proceedings of the National Academy of Sciences of the United States of America, (1990) 87/4 (1451-1455). ISSN: 0027-8424 CODEN: PNASA6 United States
SOURCE:
COUNTRY:
DOCUMENT TYPE:
                         Journal: Article
FILE SECMENT:
                                  Microbiology
                        004
                                  Clinical Biochemistry
                        029
                        English
I ANCHACE:
SUMMARY LANGUAGE:
                        English
     A number of single- and double-base substitutions have been introduced into either the polylinker region or the lacZ gene in the plasmid vector pUC19. The efficiencies of these changes upon transfection of TG-1
      bacterial cells were generally 70-80%. A strategy has been devised by which the wild-type DNA can be selectively destroyed. It is primarily based on the resistance of phosphorothioate internucleotide linkages to
      some restriction enzymes. A mismatch oligonucleotide is
      introduced into a gapped region and the gap is filled using three deoxynucleoside 5'-triphosphates and one deoxynucleoside
      5'-[.alpha.-thio]triphosphate. Reaction with a restriction enzyme that is
      unable to hydrolyze phosphorothioates ensures that the DNA containing the
      mismatch oligonucleotide is only nicked. Concomitantly, the DNA
      that does not contain the desired mutation is linearized. Subsequent
      reactions with an exonuclease and DNA polymerase I yield mutant homoduplex
      DNA for transfection.
     Medical Descriptors:
        "plasmid
      *site directed mutagenesis
      genetic engineering
      nonhuman
      article
      priority journal
      Drug Descriptors:
      *phosphorothioic acid
      (phosphorothioic acid) 10101-88-9, 13598-51-1,
      15181-41-6
L75 ANSWER 5 OF 17 EMBASE COPYRIGHT 2003 ELSEVIER INC. ALL RIGHTS RESERVED.
on STN
ACCESSION NUMBER:
                        90371788 FMRASE
DOCUMENT NUMBER:
                        1990371788
                        Chemical and enzymatic ligation of 5'-
TITLE:
                         thiophosphates of oligodeoxyribonucleotides
AUTHOR:
                        Oshevskii S.I.
                        Institute of Cytology and Genetics, Siberian Branch of the
CORPORATE SOURCE:
```

Academy of Sciences of the USSR, Novosibirsk, Russia

```
Doklady Biochemistry, (1990) 310/1-6 (15-18).
ISSN: 0012-4958 CODEN: DBIOAM
SOURCE:
COUNTRY:
                          United States
DOCUMENT TYPE:
                          Journal; Article
                                    Clinical Biochemistry
FILE SEGMENT:
                          029
                          English
LANGUAGE:
     Medical Descriptors:
         bacteriophage t4
      article
      Drug Descriptors:
      *dna
         *oligonucleotide
      *rna
      (dna) 9007-49-2; (rna) 63231-63-0
-> d ibib abs hitrn 6
L75 ANSWER 6 OF 17 HCAPLUS COPYRIGHT 2003 ACS on STN
ACCESSION NUMBER:
                                 2003:461948 HCAPLUS
                                 139:225986
DOCUMENT NUMBER:
TITLE:
                                 Comparison of different antisense strategies in
                                 mammalian cells using locked nucleic acids,
                                 2'-O-methyl RNA, phosphorothicates and small
                                 interfering RNA
AUTHOR(S):
                                 Gruenweller, Arnold; Wyszko, Eliza; Bieber, Birgit;
                                Jahnel, Ricarda; Erdmann, Volker A.; Kurreck, Jens Institut fuer Chemie-Biochemie, Freie Universitaet Berlin, Berlin, D-14195 (Germany Nucleic Acids Research ((2003), 31(12), 3185-3193 CODEN: NARHAD; ISSN: 0305=1048 Oxford University Press
CORPORATE SOURCE:
SOURCE:
PUBLISHER:
DOCUMENT TYPE:
                                 Journal
LANGUAGE:
                                English
      Locked nucleic acids (LNAs) and double-stranded small interfering RNAs
      (siRNAs) are rather new promising antisense mols. for cell culture and in
      vivo applications. Here, we compare LNA-DNA-LNA gapmer oligonucleotides and siRNAs with a phosphorothioate and a chimeric 2'-O-Me RNA-DNA gapmer with respect to their capacities to knock down the expression of the vanilloid receptor subtype 1 (VR1). LNA-DNA-LNA gapmers with four or five LNAs on either side and a central
      stretch of 10 or 8 DNA monomers in the center were found to be active gapmers that inhibit gene expression. A comparative co-transfection study showed that siRNA is the most potent inhibitor of VR1-green fluorescent
      protein (GFP) expression. A specific inhibition was obsd. with an estd.
      ICSO of 0.06 nM. An LNA gapmer was found to be the most efficient single-stranded antisense oligonucleotide.
      with an IC50 of 0.4 nM being 175-fold lower than that of commonly used
      phosphorothioates (ICSO .apprx.70 nM). In contrast, the efficiency of a 2'-O-methyl-modified oligonucleotide
      (IC50.apprx.220 nM) was 3-fold lower compared with the
      phosphorothicate. The high potency of siRNAs and chimeric LNA-DNA
      oligonucleotides make them valuable candidates for cell culture
      and in vivo applications targeting the VR1 mRNA. 15181-41-6. Phosphorothioate
      RL: BUU (Biological use, unclassified); BIOL (Biological study); USES
      (Uses)
          (RNA; gene silencing using locked nucleic acids, 2'-O-Me RNA,
          phosphorothioates and siRNA)
REFERENCE COUNT:
                                        THERE ARE 44 CITED REFERENCES AVAILABLE FOR THIS
                                44
                                       · RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
=> d ibib abs hitrn 7
L75 ANSWER 7 OF 17 HCAPLUS COPYRIGHT 2003 ACS on STN
ACCESSION NUMBER:
                                2003:339079 HCAPLUS
DOCUMENT NUMBER:
                                 139:1495
TITLE:
                                 Antisense technologies. Improvement through novel
                                 chemical modifications
AUTHOR(S):
                                 Kurreck, Jens
CORPORATE SOURCE:
                                Institut fur Chemie-Biochemie, Freie Universitat
                                Berlin, Berlin, 14195, Germany
                                 European Journal of Biochemistry (2003)
SOURCE:
                                                                                       270(8),
                                 1628-1644
                                 CODEN: EJBCAI; ISSN: 0014-2956
PUBLISHER:
                                Blackwell Publishing Ltd.
DOCUMENT TYPE:
                                 Journal; General Review
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A review. Antisense agents are valuable tools to inhibit the expression
       of a target gene in a sequence-specific manner, and may be used for
       functional genomics, target validation and therapeutic purposes. Three types of anti-mRNA strategies can be distinguished. Firstly, the use of
       single stranded antisense-oligonucleotides;
       secondly, the triggering of RNA cleavage through catalytically active oligonucleotides referred to as ribozymes; and thirdly, RNA interference induced by small interfering RNA mols. Despite the seemingly
       simple idea to reduce translation by oligonucleotides
       complementary to an mRNA, several problems have to be overcome for successful application. Accessible sites of the target RNA for oligonucleotide binding have to be identified, antisense agents
       have to be protected against nucleolytic attack, and their cellular uptake
       and correct intracellular localization have to be achieved. Major
       disadvantages of commonly used phosphorothicate DNA
       oligonuclectides are their low affinity towards target RNA mols. and their toxic side-effects. Some of these problems have been solved in "second generation" nucleotides with alkyl modifications at the 2'
       position of the ribose. In recent years valuable progress has been
       achieved through the development of novel chem. modified nucleotides with
      -improved properties such as enhanced serum stability, higher target
       affinity and low toxicity. In addn., RNA-cleaving ribozymes and
       deoxyribozymes, and the use of 21-mer double-stranded RNA mols. for RNA
       interference applications in mammalian cells offer highly efficient
       strategies to suppress the expression of a specific gene.
       15181-41-6, Phosphorothicate
       RL: BUU (Biological use, unclassified); BIOL (Biological study); USES
       (Uses)
           (comparison of different antisense strategy)
REFERENCE COUNT:
                                   131 THERE ARE 131 CITED REFERENCES AVAILABLE FOR
                                           THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE
-> d ibib abs hitrn 8
L75 ANSWER 8 OF 17 HCAPLUS COPYRIGHT 2003 ACS on STN
ACCESSION NUMBER:
                                   2003:609425 HCAPLUS
DOCUMENT NUMBER:
                                   139:241236
TITLE:
                                   A comparison of gene repair strategies in cell culture
                                   using a lacZ reporter system
Nickerson, H. D.; Colledge, W. H.
AUTHOR(S):
CORPORATE SOURCE:
                                   Department of Physiology, University of Cambridge,
                                   Cambridge, UK
Gene Therapy (2003), 10(18), 1584-1591
CODEN: GETHEC, ISSN: 0969-7128
Nature Publishing Group
SOURCE:
PURLISHER:
DOCUMENT TYPE:
                                   Journal
LANGUAGE:
                                   English
      Synthetic oligonucleotides and DNA fragments of less than 1
       kilobase (kb) have been shown to cause site-specific genetic alterations
      kilobase (kb) have been shown to cause site-specific genetic alterations in mammalian cells in culture and in vivo. We have used a lacZ reporter gene system to compare the efficiency of episomal and chromosomal gene repair in human embryonic kidney epithelial cells (HEK293), Chinese Hamster Ovary fibroblasts (CHOK1), human bronchial epithelial cells (16HBE), and mouse embryonic stem (E5) cells. The lacZ gene contains a G to A nucleotide change, (Glu to Lys mutation) that abrogates .beta.-galactosidase activity. We compared the efficiency of different
       gene repair methods to correct this mutation and restore
        beta.-galactosidase activity. We evaluated PCR-generated double-stranded
       DNA fragments of 0.52-1.9 kb, single-stranded DNA
      oligonuclectides of 20, 35, or 80 bases contg. internal phosphorothicate links, and a 68 base RNA:DNA
       oligonucleotide. All of the oligonucleotides and DNA
       fragments showed some gene repair ability with an episomal plasmid.
DNA fragments of 0.52 kb or greater gave the highest frequencies of
       episomal gene repair while single-stranded DNA
       oligonucleotides gave the highest frequency of chromosomal repair.
       In the context of a chromosomal target, antisense DNA
       oligonucleotides gave 5-fold higher frequencies of gene repair
       than their sense counterparts. The RNA:DNA chimeric
       oligonucleotide gave little or no gene repair on either a
       chromosomal or episomal target.
       15181-41-6, Phosphorothicate
       RL: BSU (Biological study, unclassified); BUU (Biological use, unclassified); BIOL (Biological study); USES (Uses)
           (comparison of gene repair strategies in cell culture using a lacZ
```

LANGUAGE:

English

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reporter system)
                                                THERE ARE 21 CITED REFERENCES AVAILABLE FOR THIS
REFERENCE COUNT:
                                                RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
=> d ibib abs hitrn 9
L75 ANSWER 9 OF 17 HCAPLUS COPYRIGHT 2003 ACS on STN
ACCESSION NUMBER:
                                       2002:658292 HCAPLUS
DOCUMENT NUMBER:
                                       137:196646
                                      Defined DNA sequences amplifiable with a universal primer pair for use in labeling materials for
TITLE:
                                       identification
INVENTOR(S):
                                       Brown, Tom; Thelwell, Nichola; Maxwell, Paula;
                                       Maxwell, Paul; Whiting, Paul
PATENT ASSIGNEE(S):
                                       Crime Solutions Limited, UK
                                      PCT Int. Appl., 23 pp. CODEN: PIXXD2
SOURCE:
DOCUMENT TYPE:
                                       Patent
LANGUAGE:
                                       English
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
        PATENT NO.
                                  KIND DATE
                                                                  APPLICATION NO. DATE
                                         20020829
        WO 2002066678
                                   A2
                                                                  WO 2002-G8759
                                                                                             20020220
        WO 2002066678
                                   A3
                   AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
              W:
                   LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
RW: GH, CM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, FT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, CQ, GW, ML, MR, NE, SN, TD, TG PRIORITY APPLN. INFO:: GB 2001-4163 A 20010220
       A method of uniquely identifying an object by labeling it with a DNA
       sequence is described. The DNA sequence has a terminal region including a
       sequence is described. The DNA sequence has a terminal region including a moiety that can be used to attach it to a substrate. Adjacent to this is a sequence by which the DNA can be released from the substrate, such as a restriction enzyme cleavage site. The remainder of the DNA is the unique identifier that includes a pair of primer binding sites sepd. by a defined and unique DNA sequence. The DNA may also contain base analogs or have a modified backbone that will prevent degrdn. of the label by nucleases.
        The DNA may also be single-stranded with the
        immobilization region in the loop of a stem loop structure. The partially
       double stranded region may serve as a primer for an initial amplification.
       Amplification and sequencing of the unique sequence identifier can be used
        to demonstrate ownership.
       15181-41-60, Thiophosphate, nucleic acid conjugates
       RL: TEM (Technical or engineered material use); USES (Uses)
(for immobilization of oligonucleotide label; defined DNA
            sequences amplifiable with universal primer pair for use in labeling
            materials for identification)
⇒ d ibib abs hitrn 10
L75 ANSWER 10 OF 17 HCAPLUS COPYRIGHT 2003 ACS on STN ACCESSION NUMBER: 2002:575095 HCAPLUS
ACCESSION NUMBER:
DOCUMENT NUMBER:
                                      137:106042
TITLE:
                                      Nuclease-based method for detecting and quantitating
                                      oligonucleotides
INVENTOR(S):
                                      Yu, Zhengrong; Baker, Brenda F.; Wu, John
Isis Pharmaceuticals, Inc., USA
PATENT ASSIGNEE(S):
SOURCE:
                                      PCT Int. Appl., 48 pp. CODEN: PIXXD2
DOCUMENT TYPE:
                                      Patent
LANGUAGE:
                                      English
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
       PATENT NO.
                                  KIND DATE
                                                                  APPLICATION NO. DATE
```

WO 2002059137

A1

20020801

W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,

WO 2001-US49702 20011023

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CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, G8, GD, GE, GH,
                GM, HR, HU, ID, IL, IN, IS. JP, KE, KG, KP, KR, KZ, LC, LK, LR,
                LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT,
                RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US,
           UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, T), TM
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
                DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF;
               BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
547 Al 20030827 EP 2001-994359 20011023
           R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
                IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
.N. INFO.: US 2000-705587
PRIORITY APPLN. INFO.:
                                                  WO 2001-US49702 W 20011023
     The invention concerns a method for quantitating an
      oligonucleotide in a sample of bodily fluid and/or ext. is
      provided. The method comprises contacting an oligonucleotide
      with a probe comprising a detectable marker and a binding moiety; placing
      the fluid or ext. in contact with a solid support to which a binding
      partner of the binding moiety is attached; contacting the fluid or ext.
      with a single-strand specific nuclease to degrade
      probe which is not hybridized to the oligonucleotide; and
      detecting a label assocd. with the marker. The method provides or the detection and/or localization of oligonucleotides, including
      administered modified oligonucleotides, for therapeutic and/or
      pharmacokinetic purposes.
      15181-41-6, Phosphorothicate
      RL: PRP (Properties)
          (nuclease-based method for detecting and quantitating
          oligonucleotides)
REFERENCE COUNT:
                                       THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS
                                       RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
-> d ibib abs hitrn 11
175 ANSWER 11 OF 17 HCAPLUS COPYRIGHT 2003 ACS on STN
                               2002:522052 HCAPLUS
ACCESSION NUMBER:
DOCUMENT NUMBER:
                               137:89420
TITLE:
                               Single-stranded circular oligonucleotide probes for
                               detection of polymorphisms in nucleic acids by
                               rolling-circle amplification (RCA)
INVENTOR(S):
                               Bandaru, Rajanikanth; Kumar, Gyanendra
PATENT ASSIGNEE(S):
                               Molecular Staging, Inc., USA
                               PCT Int. Appl., 90 pp.
CODEN: PIXXD2
SOURCE:
DOCUMENT TYPE:
                               Patent
LANGUAGE:
                               English
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
      PATENT NO.
                           KIND DATE
                                                      APPLICATION NO. DATE
      WO 2002053780
                                  (20020/11
                                                      WO 2002-US5
                                                                           20020104
                            AZ
                                  20030522
      WD 2002053780
                            A3
              AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LY, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH.
           W:
           PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR,
      BF, B), CF, CG, CI, CM, GA, GN, CQ, CW, ML, MR, NE, US 2003044794 A1 20030306 US 2001-910372 2001
      US 6635425
                            B2
                                  20031021
          EP 1347988
                                                  US 2003-465759 20030619
US 2001-259918P P 20010105
      US 2003207323
                            A1
                                  20031106
PRIORITY APPLN. INFO.:
                                                  US 2001-910372 A 20010720
                                                  WO 2002-USS
                                                                       W 20020104
     The present invention provides a novel method for ligation of
      oligonucleotides contg. 5'-phosphorothioates on complementary templates by
the action of DNA ligases. This reaction is readily applied to the
      synthesis of a single stranded circular DNA contg, a phosphorothicate directed ligation reaction by ATP dependent DNA ligase reaction is similar
      to conventional 5'-phosphate ligation. The utility of enzymic ligation in
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probing specific sequences of DNA is also described. The present invention also provides a novel non-enzymic ligation of 5'-phosphorothioates that has been applied to the synthesis of single strand phosphorothioate and phosphate circular DNA. A process for detecting the presence of a mismatch in an otherwise complementary pair of oligonucleotides is disclosed using an enzyme-based technique which shows the presence of a mismatch by failing to form a ligated single stranded DNA circle that can optionally be amplified using std. methods of rolling circle amplification.

=> d ind 11

Glass, uses

ANSWER 11 OF 17 HCAPLUS COPYRIGHT 2003 ACS on STN ICM C12Q001-68 cc 3-1 (Biochemical Genetics) Section cross-reference(s): 13 genotyping SNP single nucleotide polymorphism DNA high throughput assay; human genomic DNA SNP genotyping rolling circle amplification method; oligonucleotide rolling circle amplification nucleic acid Thermus thermophilus (DNA ligase from; single-stranded circular oligonucleotide probes for detection of polymorphisms in nucleic acids by rolling-circle amplification (RCA)) Escherichia coli Rhodothermus marinus Thermus scotoductus (DNA ligase; single-stranded circular oligonucleotide probes for detection of polymorphisms in nucleic acids by rolling-circle amplification (RCA)) Bacillus phage .phi.29 Coliphage T4 Coliphage T7 (DNA polymerase; single-stranded circular oligonucleotide probes for detection of polymorphisms in nucleic acids by rolling-circle amplification (RCA)) Primers (nucleic acid) RL: ARG (Analytical reagent use); DGN (Diagnostic use); PRP (Properties); ANST (Analytical study); BIOL (Biological study); USES (Uses) (DNA, Amplifluor, fluorescent labeled; single-stranded circular oligonucleotide probes for detection of polymorphisms in nucleic acids by rolling-circle amplification (RCA)) TT Genome (DNA; single-stranded circular oligonucleotide probes for detection of polymorphisms in nucleic acids by rolling-circle amplification (RCA)) п (biallelic SNPs; single-stranded circular oligonucleotide probes for detection of polymorphisms in nucleic acids by rolling-circle amplification (RCA)) IT rna RL: BUU (Biological use, unclassified); BIOL (Biological study); USES (Uses) (bridging oligonucleotides contg.; single-stranded circular oligonucleotide probes for detection of polymorphisms in nucleic acids by rolling-circle amplification (RCA)) Peptides, biological studies Primers (nucleic acid) Proteins RL: BUU (Biological use, unclassified); BIOL (Biological study); USES (Uses) (closed circle oligonucleotides conjugates to; single-stranded circular oligonucleotide probes for detection of polymorphisms in nucleic acids by rolling-circle amplification (RCA)) П Human (genomic DNA polymorphisms; single-stranded circular oligonucleotide probes for detection of polymorphisms in nucleic acids by rolling-circle amplification (RCA)) Conformation (hairpin loop, in oligonucleotide; single-stranded circular oligonucleotide probes for detection of polymorphisms in nucleic acids by rolling-circle amplification (RCA)) Enzymes, biological studies RL: BUU (Biological use, unclassified); BIOL (Biological study); USES (mRNA-capping, single-stranded circular oligonucleotides synthesis using; single-stranded circular oligonucleotide probes for detection of polymorphisms in nucleic acids by rolling-circle amplification (RCA))

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Plastics, uses
      RL: DEV (Device component use); USES (Uses)
         (oligonucleotide attached to solid support contg.; single-stranded circular oligonucleotide probes for detection of polymorphisms in nucleic acids by rolling-circle amplification (RCA))
     Deoxyribonucleotides
      RL: BUU (Biological use, unclassified); BIOL (Biological study); USES
      (Uses)
         (open circle oligonucleotides and bridging oligonucleotides contg.;
         single-stranded circular oligonucleotide probes for detection of
         polymorphisms in nucleic acids by rolling-circle amplification (RCA))
     DNA
      RL: ARG (Analytical reagent use); DGN (Diagnostic use); PRP (Properties);
     ANST (Analytical study); BIOL (Biological study); USES (Uses) (primer, Amplifluor, fluorescent labeled; single-
         stranded circular oligonucleotide probes for detection of
         polymorphisms in nucleic acids by rolling-circle amplification (RCA))
     Nucleic acid amplification (method)
         (rolling circle amplification; single-stranded circular oligonucleotide
         probes for detection of polymorphisms in nucleic acids by rolling-circle amplification (RCA))
     Genetic polymorphism
         (single nucleotide; single-stranded circular oligonucleotide probes for
         detection of polymorphisms in nucleic acids by rolling-circle
         amplification (RCA))
     Genotyping (method)
Nucleic acid hybridization
         (single-stranded circular oligonucleotide probes for detection of
         polymorphisms in nucleic acids by rolling-circle amplification (RCA))
      RL: ANT (Analyte); DCN (Diagnostic use); ANST (Analytical study); BIOL
      (Biological study); USES (Uses)
(single-stranded circular oligonucleotide probes
         for detection of polymorphisms in nucleic acids by rolling-circle
         amplification (RCA))
     Probes (nucleic acid)
      RL: ARG (Analytical reagent use); DGN (Diagnostic use); PRP (Properties);
      ANST (Analytical study); BIOL (Biological study); USES (Uses)
         (single-stranded circular oligonucleotide probes for detection of
         polymorphisms in nucleic acids by rolling-circle amplification (RCA))
     Oligonucleotides
      RL: BPN (Biosynthetic preparation); BSU (Biological study, unclassified);
      BIOL (Biological study); PREP (Preparation)
(single-stranded circular, bridging, synthesis of; single-stranded circular oligonucleotide probes for detection of polymorphisms in
         nucleic acids by rolling-circle amplification (RCA))
     Phosphorothicate oligonucleotides
      RL: BPN (Biosynthetic preparation); BSU (Biological study, unclassified);
     BIOL (Biological study); PREP (Preparation)
         (single-stranded circular, synthesis of; single-stranded circular
         oligonucleotide probes for detection of polymorphisms in nucleic acids
         by rolling-circle amplification (RCA))
     9015-85-4, DNA ligase
      RL: ARG (Analytical reagent use); BUU (Biological use, unclassified); DGN
      (Diagnostic use); ANST (Analytical study); BIOL (Biological study); USES
      (Uses)
         (E. coli, Thermus, Rhodothermus marinus, T4, single-stranded circular
         oligonucleotides synthesis using; single-stranded circular
         oligonucleotide probes for detection of polymorphisms in nucleic acids
         by rolling-circle amplification (RCA))
     9012-90-2D, DNA polymerase, Klenow fragment
      RL: ARG (Analytical reagent use); BUU (Biological use, unclassified); DGN
      (Diagnostic use); ANST (Analytical study); BIOL (Biological study); USES
      (Uses)
         (E. coli, phage T4 or T7, .phi.29, rolling circle amplification using; single-stranded circular oligonucleotide probes
         for detection of polymorphisms in nucleic acids by rolling-circle amplification (RCA))
     56-65-5, ATP, biological studies
RL: BUU (Biological use, unclassified); BIOL (Biological study); USES
IT
      (Uses)
         (as DNA ligase cofactor; single-stranded circular oligonucleotide
         probes for detection of polymorphisms in nucleic acids by rolling-circle amplification (RCA)
     7786-30-3, Magnesium chloride (MgCl2), biological studies RL: BUU (Biological use, unclassified); BIOL (Biological study); USES
      (Uses)
         (in ligation reaction buffer; single-stranded circular oligonucleotide
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probes for detection of polymorphisms in nucleic acids by
    rolling-circle amplification (RCA))
25952-53-8, EDC
TT
     RL: BUU (Biological use, unclassified); BIOL (Biological study); USES
     (Uses)
        (in single-stranded circular oligonucleotide synthesis; single-stranded
        circular oligonucleotide probes for detection of polymorphisms in
        nucleic acids by rolling-circle amplification (RCA))
     9037-46-1, Exonuclease I
     RL: BUU (Biological use, unclassified); BIOL (Biological study); USES
     (Uses)
        (ligation reaction products treated with; single-stranded circular
        oligonucleotide probes for detection of polymorphisms in nucleic acids
        by rolling-circle amplification (RCA))
    7704-34-9, Sulphur, biological studies
RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (of 5'phosphorothioate group not used as bridging atom for
        single-stranded circular oligonucleotide synthesis; circular
        oligonucleotide probes for detection of polymorphisms in nucleic acids .
     by rolling-circle amplification (RCA))
9012-90-2, Taq DNA ligase 37259-52-2, Ampligase
     RL: ARG (Analytical reagent use); BUU (Biological use, unclassified); DGN
     (Diagnostic use); ANST (Analytical study); BIOL (Biological study); USES
     (Uses)
        (single-stranded circular oligonucleotides synthesis using:
        single-stranded circular oligonucleotide probes for detection of
        polymorphisms in nucleic acids by rolling-circle amplification (RCA))
                   440688-21-1 440688-22-2 440688-23-3 440688-24-4, 5:
     PN: WOO2053780 SEQID: 5 unclaimed DNA 440688-25-5, 6: PN: WOO2053780
                              440688-26-6, 7: PN: W002053780 SEQID: 7 unclaimed
     SEQID: 6 unclaimed DNA
     DNA 440688-27-7, 8: PN: WOO2053780 SEQID: 8 unclaimed DNA 440688-28-8
                   440688-30-2
                                  440688-31-3
                                                440688-32-4
                                                               440688-33-5
     440688-29-9
                   440688-35-7
                                  440688-36-8
                                                440688-37-9
                                                                440688-38-0
     440688-34-6
                   440688-40-4
     440688-39-1
     RL: PRP (Properties)
        (unclaimed nucleotide sequence; single-stranded circular
        oligonucleotide probes for detection of polymorphisms in nucleic acids
        by rolling-circle amplification (RCA))
-> d ibib abs hitrn 12
    ANSWER 12 OF 17 HCAPLUS COPYRIGHT 2003 ACS on STN
                         2002:90226 HCAPLUS
ACCESSION NUMBER:
DOCUMENT NUMBER:
                          136:145278
TITLE:
                          Use of modified oligonucleotide to down-regulate gene
                          expression
                          Agrawal, Sudhir; Diasio, Robert B.; Zhang, Zhang
INVENTOR(S):
PATENT ASSIGNEE(S):
                          Hybridon, Inc., USA
                          PCT Int. Appl., 71 pp.
SOURCE:
                          CODEN: PIXXD2
DOCUMENT TYPE:
                          Patent
LANGUAGE:
                          English
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
     PATENT NO.
                      KIND DATE
                                             APPLICATION NO. DATE
     WO 2002008420
                       A2
                             /200202/31
                                             WD 2001-US18338 20010606
     WO 2002008420
                       A3
                            (20021017
            SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
             GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
             DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG

B1 20030819 US 2000-587934 20000606
     US 6608035
PRIORITY APPLN. INFO.:
                                             2000-587934
                                                              20000606
                                          US 1994-328520
                                                           A2 19941025
                                          US 1996-709910
                                                           B2 19960909
                                         US 1996-758005
                                                           B1 19961127
AB
    Disclosed is a method of down-regulating the expression of a gene in an
     animal, wherein a pharmacol. formulation comprising a chimeric
     oligonucleotide complementary to the gene is orally administered to an
     animal. The oligonucleotide administered has at least one
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phosphorothioate internucleotide linkage and at least one
     alkylphosphonate, phosphorodithioate, alkylphosphonothioate,
     phosphoramidate, phosphoramidite, phosphate ester, carbamate, carbonate,
     phosphate triester, acetamidate, or carboxymethyl ester internucleotide
     15181-41-6, Phosphorothioate
     RL: ARG (Analytical reagent use); BSU (Biological study, unclassified); ANST (Analytical study); BIOL (Biological study); USES (Uses)
        (internucleoside linkage; use of modified oligonucleotide to
        down-regulate gene expression)
=> d ind 12
L75 ANSWER 12 OF 17 HCAPLUS COPYRIGHT 2003 ACS on STN
     IOH C12N015-11
     ICS CO7H021-00; A61K031-7125; A61P025-28; A61P031-00; A61P033-00
     1-12 (Pharmacology)
     Section cross-reference(s): 3, 14
     modified oligonucleotide drug gene expression regulation
     Lymphoma
        (Burkitt's; use of modified oligonucleotide to down-regulate gene
        expression)
IT
     Trypanosoma cruzi
        (Chagas' disease from; use of modified oligonucleotide to down-regulate
        gene expression)
IT
     Leukemia
        (T-cell,
                , adult; use of modified oligonucleotide to down-regulate gene
        expression)
     Oligonucleotides
     RL: BSU (Biological study, unclassified); SPN (Synthetic preparation); THU
     (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
        (acetamidate linked; use of modified oligonucleotide to down-regulate
        gene expression)
П
     Ameba
        (amebiasis; use of modified oligonucleotide to down-regulate gene
        expression)
П
     Gene
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
        (cellular, oligonucleotide is complementary to; use of modified
        oligonucleotide to down-regulate gene expression)
TT
     Disease, animal
        (cryptoporidiosis, trichomoniasis; use of modified oligonucleotide to
        down-regulate gene expression)
IT
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
(expression; use of modified aligonucleotide to down-regulate gene
        expression)
\mathbf{\Pi}
     Filaria
        (filariasis; use of modified oligonucleotide to down-regulate gene
        expression)
π
     Disease, animal
        (foot-and-mouth disease; use of modified oligonucleotide to
        down-regulate gene expression)
IT
     Pathogen
     Virus
        (gene. oligonucleotide is complementary to: use of modified
        oligonucleotide to down-regulate gene expression)
     Intestine, disease
        (giardiasis; use of modified oligonucleotide to down-regulate gene
        expression)
     Human herpesvirus 3
        (herpes zoster from; use of modified oligonucleotide to down-regulate
        gene expression)
IT
     Ascarid
        (infestation with, Ascariasis; use of modified oligonucleotide to
        down-regulate gene expression)
     Pharynx, neoplasm
         (nasopharynx, carcinoma; use of modified oligonucleotide to
        down-regulate gene expression)
п
     Human herpesvirus
        (oral and genital; use of modified oligonucleotide to down-regulate
        gene expression)
П
     Drug delivery systems
        (oral; use of modified oligonucleotide to down-regulate gene
        expression)
TT
     Wart
```

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(papilloma; use of modified oligonucleotide to down-regulate gene
     Oligonucleotides
      RL: BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
          (phosphoramidite linked; use of modified oligonucleotide to
          down-regulate gene expression)
          (schistosomiasis from; use of modified oligonucleotide to down-regulate
         gene expression)
IT
     Toxoplasma gondii
          (toxoplasmosis from; use of modified oligonucleotide to down-regulate
          gene expression)
     AIDS (disease)
      Alzheimer's disease
      Blood plasma
      Drug metabolism
      Hepatitis
      Influenza
      Malaria
      Mammalia
      Parasite
      Pneumocystis
      Trichinella
      Trichomonacides
         (use of modified oligonucleotide to down-regulate gene expression)
      Proteins
      RL: BSU (Biological study, unclassified); BIOL (Biological study)
          (use of modified oligonucleotide to down-regulate gene expression)
     Oligonucleotides
        Phosphorothicate oligonucleotides
      RL: BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation);
      USES (Uses)
         (use of modified oligonucleotide to down-regulate gene expression)
П
     Human herpesvirus 3
          (varicella from: use of modified oligonucleotide to down-regulate gene
          expression)
TT
      Papilloma
         (warts; use of modified oligonucleotide to down-regulate gene
         expression)
IT
      Fever and Hyperthermia
         (yellow; use of modified oligonucleotide to down-regulate gene
          expression)
      463-77-4, Carbamic acid, biological studies 993-13-5 3812-32-6, Carbonate, biological studies 7664-38-2D, Phosphoric acid, triesters,
      Carbonate, biological studies 7664-38-2D, Phosphoric a biological studies 13598-36-2D, Phosphonic acid, alkyl
      15181-41-6, Phosphorothioate
22638-09-1, Phosphoramidate
                                          16481-04-2, Carboxy methyl ester
      RL: ARG (Analytical reagent use); BSU (Biological study, unclassified);
      ANST (Analytical study); BIOL (Biological study); USES (Uses)
          (internucleoside linkage; use of modified oligonucleotide to
          down-regulate gene expression)
                      393599-16-1
                                       393599-17-2
      393599-15-0
                                                         393599-18-3
      393599-20-7
                      393599-21-8
                                        393599-22-9
                                                         393599-23-0
      393599-25-2
                      393599-26-3
                                        393599-27-4
                                                         393599-28-5
                                                                          393599-29-6
      RL: PRP (Properties)
          (unclaimed nucleotide sequence; use of modified oligonucleotide to
          down-regulate gene expression)
=> d ibib abs hitrn 13
L75 ANSWER 13 OF 17 HCAPLUS COPYRIGHT 2003 ACS on STN
ACCESSION NUMBER:
                              2002:457395 HCAPLUS
DOCUMENT NUMBER:
                              137:259481
                              Separation of Synthetic Oligonucleotide Dithioates
TITLE:
                              from Monothiophosphate Impurities by Anion-Exchange
                              Yang, Xianbin; Hodge, Richard P.; Luxon, Bruce A.;
Shope, Robert; Gorenstein, David G.
Sealy Center for Structural Biology and Department of
AUTHOR(S):
CORPORATE SOURCE:
                              Human Biological Chemistry & Genetics, University of Texas Medical Branch at Galveston, TX, 77555-1157, USA Analytical Biochemistry (2002), 306(1), 92-99 CODEN: ANBCA2; ISSN: 0003-2697
SOURCE:
PUBLISHER:
                              Elsevier Science
```

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DOCUMENT TYPE:
                            Journal
 LANGUAGE:
                            English
      A method using a strong anion-exchange liq.-chromatog. column, Mono-Q, has
      been developed for high-resoln. anal. and purifn. of oligonucleotide
      dithioates, which were synthesized by an automated, solid-phase,
      phosphorothioamidite chem. High-resoln. sepn. of oligonucleotide
      phosphorodithicates from monothicphosphate impurities was obtained.
      High-resoln, sepn. was also demonstrated at pH 8. The sepn. of
      oligonucleotide dithioates was found to be linearly dependent on the no.
      of sulfurs for the same sequence length. Thiocyanate, SCN-, as eluting anion, can be used to purify oligonucleotides contg. a high percentage of
      phosphorodithicate linkages in lower salt concns. and provides better
      sepn. than chloride as eluting anion.
      15181-41-6P, Phosphorothioate
      RL: BYP (Byproduct); PREP (Preparation)
          (mono-, di-; sepn. of synthetic oligonucleotide dithioates from
         monothiophosphate impurities by anion-exchange chromatog. on a mono-Q
 REFERENCE COUNT:
                                   THERE ARE 39 CITED REFERENCES AVAILABLE FOR THIS
                                   RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
 => d ind 13
 L75 ANSWER 13 OF 17 HCAPLUS COPYRIGHT 2003 ACS on STN
      9-3 (Biochemical Methods)
      Section cross-reference(s): 6
      monoQ column oligonucleotide dithioate chromatog purifn; monothiophosphate
      oligonucleotide phosphorodithioate sepn
. 11
          (8, sepn. at; sepn. of synthetic oligonucleotide dithioates from
          monothiophosphate impurities by anion-exchange chromatog. on a mono-Q
 п
      Ion exchange chromatography
          (high-performance; sepn. of synthetic oligonucleotide dithioates from
          monothiophosphate impurities by anion-exchange chromatog. on a mono-Q
 TT
      Phosphorothicate oligonuclectides
      RL: PUR (Purification or recovery); PREP (Preparation)
          (sepn. of synthetic oligonucleotide dithioates from monothiophosphate
          impurities by anion-exchange chromatog. on a mono-Q column)
 IT
      302-04-5, Thiocyanate, uses
      RL: NUU (Other use, unclassified); USES (Uses)
         (eluting anion of; sepn. of synthetic oligonucleotide dithioates from monothiophosphate impurities by anion-exchange Chromatog. on a mono-Q
         Column)
      15181-41-6P, Phosphorothioate
      RL: BYP (Byproduct); PREP (Preparation)
         (mono-, di-; sepn. of synthetic oligonucleotide dithioates from monothiophosphate impurities by anion-exchange chromatog. on a mono-Q
         column)
      131159-51-8, Mono Q HR 10/10
      RL: NUU (Other use, unclassified); USES (Uses)
          (sepn. of synthetic oligonucleotide dithioates from monothiophosphate
          impurities by anion-exchange chromatog. on a mono-Q column)
 => d ibib abs hitrn 14
 L75 ANSWER 14 OF 17 HCAPLUS COPYRIGHT 2003 ACS on STN
 ACCESSION NUMBER:
                            2001:868734 HCAPLUS
 DOCUMENT NUMBER:
                            136:1591
 TITLE:
                            Genotyping methods to detect DNA sequence
                            polymorphisms and haplotypes
 INVENTOR(S):
                            Stanton, Vincent P.,
                            Variagenics, Inc., USA
PCT Int. Appl., 166 pp.
 PATENT ASSIGNEE(S):
 SOURCE:
                            CODEN: PIXXD2
 DOCUMENT TYPE:
                            Patent
 LANGUAGE:
                            English
 FAMILY ACC. NUM. COUNT:
                            3
 PATENT INFORMATION:
      PATENT NO.
                         KIND DATE
                                               APPLICATION NO.
                                                                  DATE
                                                WO 2001-US16577 20010523
                         A2
                               20011129
      WO 2001090419
      WO 2001090419
                          A3
                               20030710
          W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
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CO, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS,
                   LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO,
                   RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
             RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,
                   BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
36 B1 20021105 US 2000-696998 20001025
       US 6475736
PRIORITY APPLN. INFO.:
                                                           US 2000-206613P P
                                                                                         20000523
                                                           US 2000-696998
                                                                                    A2 20001025
                                                           US 2000-697013
                                                                                    A2 20001025
                                                           US 2000-697028
                                                                                   A2 20001025
```

Methods for detg. genotypes and haplotypes of genes are claimed. Also described are single nucleotide polymorphisms (SNPs) and haplotypes in the ApoE gene and their use in methods of this invention. Methods of the invention involve allele enrichment methods such as allele capture, allele-specific amplification, and allele-specific restriction endonuclease digestion. Allele capture means phys. sepn. of either single-stranded or double-stranded DNA. This can be accomplished by protein or nucleic acid reagents, such as disabled restriction enzymes zinc-finger DNA-binding proteins, and covalent crosslinking agents, which have affinity for specific alleles. The captured complexes are then sepd. from the nucleic acid mixt. by reagents such as antibody-coated beads or streptavidin. Allele-specific amplification can be accomplished by strand obstruction, such as formation of stable secondary structures, or modified primers such as covalently crosslinkable primers. Lastly, allele-specific restriction methods for genotyping can be accomplished by triplex-mediated protection, primer-mediated creation of polymorphic restriction sites, and other variations, followed by amplification, direct nucleotide sequencing, or capture and size or sequence anal. Allele-specific primers were designed to det. haplotypes of nucleotide 186 T/C and 597 A/G polymorphisms in the dihydropyrimidine dehydrogenase gene. The primers are allele-specific because they induce hairpin loop formation when the "correct" nucleotide is present at the polymorphic site. The hairpin loop structure inhibits annealing of new primers and further amplification. PCR products were digested with BsrDI restriction endonuclease and analyzed by agarose gel electrophoresis. A T/C SNP at genomic site 21250 in the human ApoE gene results in a cysteine to arginine substitution at position 176 of the ApoE protein. For genotyping the T/C SNP, a loop primer and reverse primer were designed to amplify the target and introduce FokI and FspI restriction enzyme cleavage sites. Digestion with FokI and FspI produced allele-specific DNA fragments which were sequenced by mass spectrometry. Fourteen polymorphic sites for the ApoE gene and exptl. derived haplotypes for some or all of these polymorphisms are provided.

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=> d ind 14
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ANSWER 14 OF 17 HCAPLUS COPYRIGHT 2003 ACS on STN
L75
      ICM C120001-68
      3-1 (Biochemical Genetics)
cc
      Section cross-reference(s): 9, 13
      genotyping polymorphism haplotype allele DNA binding complex restriction endonuclease; human gene ApoE SNP genotype haplotype PCR sequence analysis
ST
      Gene, animal
      RL: ANT (Analyte); BUU (Biological use, unclassified); ANST (Analytical study); BIOL (Biological study); USES (Uses)
(APOE; genotyping methods to detect DNA sequence polymorphisms and
          haplotypes)
      Quaternary structure
          (DNA triplex, allele-specific; genotyping methods to detect DNA
          sequence polymorphisms and haplotypes)
      RL: ARG (Analytical reagent use); BUU (Biological use, unclassified); ANST
      (Analytical study); BIOL (Biological study); USES (Uses)
          (DNA-binding; genotyping methods to detect DNA sequence polymorphisms
          and haplotypes)
      Primers (nucleic acid)
RL: ARG (Analytical reagent use); BUU (Biological use, unclassified); ANST
      (Analytical study); BIOL (Biological study); USES (Uses)
          (DNA; genotyping methods to detect DNA sequence polymorphisms and
          haplotypes)
      Enzymes, biological studies
      RL: ARG (Analytical reagent use); BUU (Biological use, unclassified); ANST (Analytical study); BIOL (Biological study); USES (Uses)
          (RecA; genotyping methods to detect DNA sequence polymorphisms and
```

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haplotypes)
IŦ
     Molecular association
         (allele-specific DNA-binding; genotyping methods to detect DNA sequence
         polymorphisms and haplotypes)
\Pi
     Hydrogen bond
        (allele-specific; genotyping methods to detect DNA sequence polymorphisms and haplotypes)
TT
     RNA
     RL: ARG (Analytical reagent use); BUU (Biological use, unclassified); ANST
     (Analytical study); BIOL (Biological study); USES (Uses)
         (aptamer: genotyping methods to detect DNA sequence polymorphisms and
        haplotypes)
     Peptide nucleic acids
     Proteins
     Transcription factors
     RL: ARG (Analytical reagent use); BUU (Biological use, unclassified); ANST
     (Analytical study); BIOL (Biological study); USES (Uses)
(biotinylated or immobilized; genotyping methods to detect DNA sequence
        polymorphisms and haplotypes)
IT
     ONA
     RL: ANT (Analyte); BUU (Biological use, unclassified); ANST (Analytical
     study); BIOL (Biological study); USES (Uses)
        (double-stranded; genotyping methods to detect DNA sequence
        polymorphisms and haplotypes)
     Alleles
     Crosslinking
     Genotypes
     Genotyping (method)
     Immunoassay
     Nucleic acid amplification (method)
     PCR (polymerase chain reaction)
     RFLP (restriction fragment length polymorphism)
         (genotyping methods to detect DNA sequence polymorphisms and
        haplotypes)
П
     Gene, animal
     cDNA
     RL: ANT (Analyte); BUU (Biological use, unclassified); ANST (Analytical
     study); BIOL (Biological study); USES (Uses)
         (genotyping methods to detect DNA sequence polymorphisms and
        haplotypes)
IT
     Oligonucleotides
     RL: ARG (Analytical reagent use); BUU (Biological use, unclassified); ANST
     (Analytical study); BIOL (Biological study); USES (Uses)
         (genotyping methods to detect DNA sequence polymorphisms and
         haplotypes)
П
     Peptide nucleic acids
     RL: ARG (Analytical reagent use); BUU (Biological use, unclassified); ANST (Analytical study); BIOL (Biological study); USES (Uses)
         (genotyping methods to detect DNA sequence polymorphisms and
        haplotypes)
п
     Phosphorothicate oligonucleotides
     RL: ARG (Analytical reagent use); BUU (Biological use, unclassified); ANST (Analytical study); BIOL (Biological study); USES (Uses)
         (genotyping methods to detect DNA sequence polymorphisms and
        haplotypes)
     Primers (nucleic acid)
     RL: ARG (Analytical reagent use); BUU (Biological use, unclassified); ANST
     (Analytical study); BIOL (Biological study); USES (Uses)
         (genotyping methods to detect DNA sequence polymorphisms and
        haplotypes)
     Proteins
     RL: ARG (Analytical reagent use); BUU (Biological use, unclassified); ANST
     (Analytical study); BIOL (Biological study); USES (Uses)
         (genotyping methods to detect DNA sequence polymorphisms and
        haplotypes)
     Transcription factors
     RL: ARG (Analytical reagent use); BUU (Biological use, unclassified); ANST
     (Analytical study); BIOL (Biological study); USES (Uses)
         (genotyping methods to detect DNA sequence polymorphisms and
        haplotypes)
     Peptides, biological studies
     RL: ARG (Analytical reagent use); BUU (Biological use, unclassified); ANST (Analytical study); BIOL (Biological study); USES (Uses)
         (histidine-contg., ligand tag; genotyping methods to detect DNA
         sequence polymorphisms and haplotypes)
     Oligonucleotides
     RL: ARG (Analytical reagent use); BUU (Biological use, unclassified); ANST
     (Analytical study); BIOL (Biological study); USES (Uses)
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(immobilized; genotyping methods to detect DNA sequence polymorphisms
         and haplotypes)
     Oligonucleotides
IT
     RL: ARG (Analytical reagent use); BUU (Biological use, unclassified); ANST
     (Analytical study); BIOL (Biological study); USES (Uses)
         (labeled, biotinylated; genotyping methods to detect DNA sequence
         polymorphisms and haplotypes)
     Magnetic particles
         (ligand tag; genotyping methods to detect DNA sequence polymorphisms
         and haplotypes)
IT
     Antibodies
     Avidins
     RL: ARG (Analytical reagent use); BUU (Biological use, unclassified); ANST.
     (Analytical study); BIOL (Biological study); USES (Uses)
         (ligand tag; genotyping methods to detect DNA sequence polymorphisms
         and haplotypes)
П
     Conformation
         (loop, nucleic acid, D-loop, allele-specific; genotyping methods to
         detect DNA sequence polymorphisms and haplotypes)
п
     DNA sequence analysis
         (mass spectrometric; genotyping methods to detect DNA sequence
         polymorphisms and haplotypes)
     Nucleic acid bases
     RL: BUU (Biological use, unclassified); BIOL (Biological study); USES
         (mass-modified; genotyping methods to detect DNA sequence polymorphisms
        and haplotypes)
     Imaging
П
         (optical mapping; genotyping methods to detect DNA sequence
         polymorphisms and haplotypes)
TT
     Nucleic acid bases
     RL: BSU (Biological study, unclassified); BIOL (Biological study)
(pairing, allele-specific; genotyping methods to detect DNA sequence
        polymorphisms and haplotypes)
IT
     DNA
     RL: ARG (Analytical reagent use); BUU (Biological use, unclassified); ANST
     (Analytical study); BIOL (Biological study); USES (Uses)
         (primer; genotyping methods to detect DNA sequence polymorphisms and
        haplotypes)
     Genetic element
     RL: BUU (Biological use, unclassified); BIOL (Biological study); USES
     (Uses)
         (restriction endonuclease cleavage site; genotyping methods to detect
        DNA sequence polymorphisms and haplotypes)
     Polyamides, biological studies
     RL: ARG (Analytical reagent use); BUU (Biological use, unclassified); ANST (Analytical study); BIOL (Biological study); USES (Uses) (sequence-specific DNA-binding; genotyping methods to detect DNA sequence polymorphisms and haplotypes)
TT
     Genetic polymorphism
         (single nucleotide; genotyping methods to detect DNA sequence
        polymorphisms and haplotypes)
     RL: ANT (Analyte); BUU (Biological use, unclassified); ANST (Analytical
     study); BIOL (Biological study); USES (Uses)
(single-stranded; genotyping methods to detect DNA
        sequence polymorphisms and haplotypes)
     Separation
        (size selection; genotyping methods to detect DNA sequence
        polymorphisms and haplotypes)
IT
     Immunoassav
         (solid-phase; genotyping methods to detect DNA sequence polymorphisms
        and haplotypes)
     Proteins
     RL: ARG (Analytical reagent use); BUU (Biological use, unclassified); ANST
     (Analytical study); BIOL (Biological study); USES (Uses)
         (zinc finger-contg., biotinylated or immobilized; genotyping methods to
        detect DNA sequence polymorphisms and haplotypes)
IT
     RL: ARG (Analytical reagent use); BUU (Biological use, unclassified); ANST (Analytical study); BIOL (Biological study); USES (Uses)
         (zinc finger-contg.; genotyping methods to detect DNA sequence
        polymorphisms and haplotypes)
     9012-90-2, DNA polymerase
     RL: ARG (Analytical reagent use); BUU (Biological use, unclassified); ANST
     (Analytical study); BIOL (Biological study); USES (Uses)
         (T4 and I, exonuclease; genotyping methods to detect DNA sequence
        polymorphisms and haplotypes)
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66-97-7, Psoralen 22542-10-5D, complexes, biological studies 146237-51-6 146237-52-7 146237-53-8
      RL: ARG (Analytica) reagent use); BUU (Biological use, unclassified); ANST
      (Analytical study); BIOL (Biological study); USES (Uses)
          (crosslinking agent; genotyping methods to detect DNA sequence
          polymorphisms and haplotypes)
      9026-89-5, Dihydropyrimidine dehydrogenase
      RL: BSU (Biological study, unclassified); BIOL (Biological study)
          (gene for; genotyping methods to detect DNA sequence polymorphisms and
          haplotypes)
      9037-44-9, Escherichia coli exonuclease III 9075-08-5, Restriction
      endonuclease
                        37228-74-3, Exonuclease 37367-70-7, Lambda exonuclease
      58513-62-5, Nuclease, bacteriophage T7 exodeoxyribo-
      81295-34-3, Restriction endonuclease PvuII 81458-03-9, Restriction
      endonuclease FokI 85340-94-9, Bal31 exonuclease 92228-44-9,
Restriction endonuclease NcoI 103780-20-7, NotI restriction endonuclease
      107824-63-5 135340-89-5, Restriction endonuclease N.BstNBI
      174632-11-2, Restriction endonuclease BsgI
                                                             189088-83-3, Restriction
      endonuclease BsrDI
      RL: ARG (Analytical reagent use); BUU (Biological use, unclassified); ANST
      (Analytical study); BIOL (Biological study); USES (Uses)
          (genotyping methods to detect DNA sequence polymorphisms and
          haplotypes)
                            7440-02-0, Nickel, biological studies
      58-85-5, Biotin
      Streptavidin
      RL: ARG (Analytical reagent use); BUU (Biological use, unclassified); ANST
      (Analytical study); BIOL (Biological study); USES (Uses)
          (ligand tag; genotyping methods to detect DNA sequence polymorphisms
          and haplotypes)
      9025-82-5, Phosphodiesterase
      RL: ARG (Analytical reagent use); BUU (Biological use, unclassified); ANST
      (Analytical study); BIOL (Biological study); USES (Uses)
          (snake venom type I; genotyping methods to detect DNA sequence
          polymorphisms and haplotypes)
-> d ibib abs hitrn 15
L75 ANSWER 15 OF 17 HCAPLUS COPYRIGHT 2003 ACS on STN
ACCESSION NUMBER:
                               1999:115526 HCAPLUS
DOCUMENT NUMBER:
                               130:292382
TITLE:
                               High sequence fidelity in a non-enzymic DNA
                               autoligation reaction
                              Xu, Yanzheng; Kool, Eric T.
Department of Chemistry, University of Rochester,
Rochester, NY, 14627, USA
AUTHOR($):
CORPORATE SOURCE:
                              Nucleic Acids Research (1999), 27(3), 875-881
CODEN: NARHAD; ISSN: 0305-1048
SOURCE:
                               Oxford University Press
PUBLISHER:
DOCUMENT TYPE:
                               Journal
LANGUAGE:
                              English
      The success of oligonucleotide ligation assays in probing
      specific sequences of DNA arises in large part from high enzymic selectivity against base mismatches at the ligation junction. We describe
      here a study of the effect of mismatches on a new non-enzymic, reagent-free method for ligation of oligonucleotides. In this approach, two oligonucleotides bound at adjacent sites on a
      complementary strand undergo autoligation by displacement of a 5'-end iodide with a 3'-phosphorothioate group. The data show that this ligation proceeds somewhat more slowly than ligation by T4 ligase.
      but with substantial discrimination against single base mismatches both at
      either side of the junction and a few nucleotides away within one of the oligonucleotide binding sites. Selectivities of >100-fold against a single mismatch are obsd. in the latter case. Expts. at varied concns.
      and temps. are carried out both with the autoligation of two adjacent
      linear oligonucleotides and with intramol. autoligation to yield
      circular "padlock" DNAs. Application of optimized conditions to discrimination of an H-ras codon 12 point mutation is demonstrated with a
      single-stranded short DNA target.
     15181-41-6, Phosphorothicate
      RL: BAC (Biological activity or effector, except adverse); BSU (Biological
      study, unclassified); BIOL (Biological study)
(autoligation by displacement of a 5'-end iodide with a 3'-
          phosphorothicate group; high sequence fidelity in a non-enzymic
          DNA autoligation reaction)
                                      THERE ARE 45 CITED REFERENCES AVAILABLE FOR THIS
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RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

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=> d ind 15
L7S ANSWER 15 OF 17 HCAPLUS COPYRIGHT 2003 ACS on STN CC 3-4 (Biochemical Genetics)
      Section cross-reference(s): 6, 9
     nonenzymic DNA autoligation reaction high sequence fidelity
      Codons
      RL: BSU (Biological study, unclassified); BIOL (Biological study)
         (12, application of optimized conditions to discrimination of an H-ras
         codon 12 point mutation is demonstrated; high sequence fidelity in a
         non-enzymic DNA autoligation reaction)
     RL: BAC (Biological activity or effector, except adverse); BPR (Biological process); BSU (Biological study, unclassified); BIOL (Biological study);
      PROC (Process)
         (autoligation; high sequence fidelity in a non-enzymic DNA autoligation
         reaction)
     Mutation
         (base-mismatching, ligation proceeds more slowly than ligation by T4
         ligase, but with discrimination against single base mismatches; high
         sequence fidelity in a non-enzymic DNA autoligation reaction)
     Gene. animal
      RL: BSU (8iological study, unclassified); BIOL (Biological study)
(c-Ha-ras, application of optimized conditions to discrimination of an
         H-ras codon 12 point mutation is demonstrated; high sequence fidelity
         in a non-enzymic DNA autoligation reaction)
TΥ
     DNA
      RL: BOC (Biological occurrence); BSU (Biological study, unclassified);
     BIOL (Biological study); OCCU (Occurrence)
(circular, autoligation of two adjacent linear oligonucleotides
         and with intramol. autoligation to yield circular "padlock" DNAs; high sequence fidelity in a non-enzymic DNA autoligation reaction)
     Mutation
         (point, application of optimized conditions to discrimination of an
         H-ras codon 12 point mutation is demonstrated; high sequence fidelity
         in a non-enzymic DNA autoligation reaction)
     15181-41-6, Phosphorothicate
      RL: BAC (Biological activity or effector, except adverse); BSU (Biological
      study, unclassified); BIOL (Biological study)
(autoligation by displacement of a 5'-end iodide with a 3'-
         phosphorothicate group; high sequence fidelity in a non-enzymic
         DNA autoligation reaction)
     20461-54-5, Iodide, biological studies
      RL: BPR (Biological process); BSU (Biological study, unclassified); BIOL
      (Biological study); PROC (Process)
         (autoligation by displacement of a 5'-end iodide with a 3'-
         phosphorothicate group; high sequence fidelity in a non-enzymic
         DNA autoligation reaction)
-> d ibib abs hitrn 16
L75 ANSWER 16 OF 17 HCAPLUS COPYRIGHT 2003 ACS on STN
                             1994:623647
ACCESSION NUMBER:
                                           HCAPLUS
DOCUMENT NUMBER:
                             121:223647
                             Enzymic preparation of single-stranded DNA containing
TITLE:
                             nuclease-resistant modified nucleotides using
                             phosphorothioate-containing primers
                             Nikiforov, Theo; Knapp, Michael R.
Molecular Tool, Inc., USA
PCT Int. Appl., 57 pp.
INVENTOR(S):
PATENT ASSIGNEE(S):
SOURCE:
                             CODEN: PIXXD2
DOCUMENT TYPE:
                             Patent
                             English
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
      PATENT NO.
                          KIND DATE
                                                  APPLICATION NO. DATE
     WO 9416090
                           A1
                                19940721
                                                  WO 1994-US771
                                                                      19940118
              AT. AU, BB, BG, BK, BY, CA, CH, CZ, DE, DK, ES, FI, GB, HU, JP, KP, KR, KZ, LK, LU. MG. MN, MW, NL, NO. NZ, PL, PT, RO, RU, SD,
               SE, SK, UA, US, VN.
          RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE,
               BF, BJ, CF, CG, CI. CM, GA, GN, ML, MR, NE, SN, TD, TG
900 A 19960521 US 1993-155746 19931123
      US 5518900
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A1

AU 9461262

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AU 1994-61262

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19961212
      AU 674211
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      EP 679190
                                                         EP 1994-907855 19940118
                              A1
                                     19951102
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                                     20030502
           R: AT, BE, CH, DE, DK, ES, FR, CB, GR, IE, IT, LI, LU, MC, NL, PT, SE
      1P 08505535
                                    19960618
                                                         JP 1994-516386
                                                                               19940118
                              T2
      JP 3330946
                              R2
                                     20021007
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                                     20030515
                                                         AT 1994-907855
                                                                               19940118
PRIORITY APPLN. INFO.:
                                                     US 1993-5061
                                                                               19930115
                                                    US 1993-155746
WO 1994-US771
                                                                               19931123
                                                                           W 19940118
      A method for generating single-stranded nucleic acid mols, that contain nuclease-resistant modified nucleotides and so are resistant to
      5'.fwdarw.3'-exonucleases are described. The method involves synthesizing
      the nucleic acid by primer extension using phosphorothioate -contg. primers. A pair of primers with one of them having a phosphorothiate-rich 5'-region and the other not contg.
      phosphorothicate nucleotides are used to amplify the target sequence. The amplification products are then digested with a
      5'.fwdarw,3'-nuclease with the hydrolysis of all of the nucleic acids
      present except for the amplification products contg. the
      phosphorothiate-rich primer. These products can be used in DNA sequencing
      and in the detn. of genetic polymorphism, esp. single base polymorphisms. If the phosphorothiates are placed at the 3'-end of the primer, then any
      residual primers in the reaction can be hydrolyzed with a
      5'.fwdarw.3'-nuclease to prevent further amplification.
=> d ind.16
      ANSWER 16 OF 17 HCAPLUS COPYRIGHT 2003 ACS on STN
      ICM C12P019-34
3-1 (Biochemical Genetics)
CC
      nuclease resistant single stranded DNA
      Deoxyribonucleic acid sequence determination
      Polymerase chain reaction
          (enzymic prepn. of single-stranded DNA contg. nuclease-resistant
          modified nucleotides using phosphorothicate-contg. primers)
IT
      Genetic polymorphism
          (single base, detn. of; enzymic prepn. of single-stranded DNA contg.
nuclease-resistant modified nucleotides using phosphorothioate
           -contg. primers)
      Deoxyribonucleic acids
      RL: BUU (Biological use, unclassified); RCT (Reactant); BIOL (Biological
      (single-stranded; enzymic prepn. of single
-stranded DNA contg. nuclease-resistant modified nucleotides
using phosphorothioate-contg. primers)
      Nucleotides, biological studies
      RL: BUU (Biological use, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses)
          (oligo-, deoxyribo-, thiophosphate-
linked, primers; enzymic prepn. of single-stranded DNA contg.
nuclease-resistant modified nucleotides using phosphorothioate
           -contg. primers)
      Deoxyribonucleic acids
      RL: BPN (Biosynthetic preparation); BIOL (Biological study); PREP
      (Preparation)
          (thiophosphate-linked, single-stranded,
          nuclease resistant; enzymic prepn. of single-stranded DNA contg. nuclease-resistant modified nucleotides using
      phosphorothioate-contg. primers)
79121-99-6, 5'.fwdarw.3'-Exonuclease
      RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); BIOL (Biological study)
          (phage T6 or .lambda.; enzymic prepn. of single-stranded DNA
          contg. nuclease-resistant modified nucleotides using
          phosphorothioate-contg. primers)
=>/d ibib abs hitrn 17
L75
      ANSWER 17 OF 17 HCAPLUS COPYRIGHT 2003 ACS on STN SSION NUMBER: 1992:229075 HCAPLUS
ACCESSION NUMBER:
DOCUMENT NUMBER:
                                 116:229075
TITLE:
                                 Phosphorothioate-based site-directed
                                 mutagenesis for single-stranded
                                 vectors
AUTHOR(S):
                                 Sayers, Jon R.; Eckstein, Fritz
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KHARE 10/007,489

CORPORATE SOURCE:

Abt. Chem., Max Planck Inst. Exp. Med., Heidelberg, D-6900/1, Germany

SOURCE:

Directed Mutagen. (1991), 49-69. Editor(s): McPherson, M. J. IRL: Oxford, UK.

CODEN: 57RUAL

DOCUMENT TYPE: LANGUAGE:

Conference; General Review

English

A review with 22 refs. The phosphorothioate-based oligonucleotide-directed mutagenesis method is based on the observation that certain restriction endonucleases are incapable of hydrolyzing phosphorothicate internuclectidic linkages. Thus, double-stranded DNA contg. phosphorothicate linkages in one strand only may be nicked in the non-substituted strand. In this mutagenesis procedure the mismatch oligonucleotide primer is annealed to the (+)strand of a single-stranded circular phage DNA. The primer is extended by a polymn, reaction in which one of the natural deoxynucleoside triphosphates is replaced by the corresponding deoxynucleotide 5'-O-(1-thiotriphosphate), dNTP.alpha.S. Thus, phosphorothioate groups are incorporated exclusively into the (-)strand of the newly synthesized RF-IV DNA. This results in a strand asymmetry which may be exploited. The methods, scope, and limitations of the procedure are discussed.

15181-41-6, Phosphorothicate RL: BIOL (Biological study)

(for site-directed mutagenesis of single-stranded DNA vectors)

=> d ind 17

L75 ANSWER 17 OF 17 HCAPLUS COPYRIGHT 2003 ACS on STN CC 3-0 (Biochemical Genetics)

Section cross-reference(s): 9

mutagenesis site directed phosphorothicate review

Genetic vectors

(single-stranded DNA, site-directed phosphorothioate-based mutagenesis of)

TT Mutation

(site-specific, phosphorothioate-based, for single-stranded DNA vectors) 15181-41-6, Phosphorothioate RL: BIOL (Biological study)

(for site-directed mutagenesis of single-stranded DNA vectors)